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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY S		
09/537,180	03/29/2000	THE THAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
		Donald R. Owen	WPB40219A	6869	
	7590 12/20/2004		EXAMINER		
P.O. BOX 199	-		SAUCIER, SANDRA E		
ALEXANDRI	A, VA 22320		ART UNIT	PAPER NUMBER	
			1651		
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Please find below and/or attached an Office communication concerning this application or proceeding.

Office Action Summary Saminer Sandra Saucier Sandr		Application No.	Applicant(s)				
Examiner Sandra Saucier 1651		09/537,180					
The MALLING DATE of this communication appears on the cover sheet with the correspondence address Period for Reply A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 3 MONTH(S) FROM THE MALLING DATE of THIS COMMUNICATION. Extensions of time ray be available under the provisions of 37 CPR 1 136(a). Extensions of time ray be available under the provisions of 37 CPR 1 136(a). If the period for oney specified above, the maximum statution period will apply and will exercise fix (8) MONTH 50 (2) and see with the correspondent timely. If MO period for regly is appoiled above, the maximum statution period will apply and will exercise fix (8) MONTH 50 (2) and see will be considered intelly. If MO period for regly is appoiled above, the maximum statution period will apply and will exercise fix (8) MONTH 50 (2) and see with the correspondent timely. If MO period for regly is appoiled above, the maximum statution period will apply and will exercise fix (8) MONTH 50 (2) and the consideration. If MO period for regly is appoiled above, the maximum statution period will apply and will exercise fix (8) MONTH 50 (2) (2) and (2) (2) (2) (2) (2) (2) (2) (2) (2) (2)	Office Action Summary	Examiner					
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This action is FINAL. Zb) This action is non-final.	Extensions of time may be available under the provisions of 37 CFR 1 after SIX (6) MONTHS from the mailing date of this communication. If the period for reply specified above is less than thirty (30) days, a re If NO period for reply is specified above, the maximum statutory period. Failure to reply within the set or extended period for reply will, by statut Any reply received by the Office later than three months after the mail.	I. 1.136(a). In no event, however, may a reply within the statutory minimum of thind d will apply and will expire SIX (6) MON	reply be timely filed ty (30) days will be considered timely. ITHS from the mailing date of this communication.				
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Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213. Disposition of Claims	2a)⊠ This action is FINAL . 2b)□ This action is non-final.						
Disposition of Claims 4)	3) Since this application is in condition for allowance except for formal matters, prosecution as to the merits is						
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4a) Of the above claim(s) 268-292 is/are withdrawn from consideration. 5	Disposition of Claims						
5) Claim(s) is/are allowed. 6) Claim(s) is/are objected to. 7) Claim(s) is/are objected to. 8) Claim(s) is/are objected to. 8) Claim(s) are subject to restriction and/or election requirement. **Application Papers** 9) The specification is objected to by the Examiner. 10) The drawing(s) filed on is/are: a) accepted or b) objected to by the Examiner. Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a). Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d). 11) The oath or declaration is objected to by the Examiner. Note the attached Office Action or form PTO-152. **riority under 35 U.S.C. § 119** 12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of: 1. Certified copies of the priority documents have been received. 2. Certified copies of the priority documents have been received in Application No 3. Copies of the certified copies of the priority documents have been received in Application No 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. Acknowledgment is not received. Application (PTO-413) Paper Not/Mail Date Notice of Draftsperson's Patent Drawing Review (PTO-948) Paper Not/Mail Date Notice of Information Disclosure Statement(s) (PTO-1449 or PTO/SB/08) Internation Disclosure Statement(s) (PTO-1452) Paper Not/Mail Date Notice of Information Disclosure Statement(s) (PTO-1449 or PTO/SB/08) Paper Not/Mail Date Notice of Information Disclosure Statement(s) (PTO-1449 or PTO/SB/08) Paper Not/Mail Date	4) Claim(s) <u>224,228,229 and 232-292</u> is/are pending in the application.						
6) Claim(s) 224.228,229 and 232-267 is/are rejected. 7) Claim(s) is/are objected to. 8) Claim(s) is/are objected to. 8) Claim(s) is/are objected to. 8) Claim(s) is/are objected to restriction and/or election requirement. Application Papers 9) The specification is objected to by the Examiner. 10) The drawing(s) filed on is/are: a) accepted or b) objected to by the Examiner. Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a). Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d). 11) The oath or declaration is objected to by the Examiner. Note the attached Office Action or form PTO-152. riority under 35 U.S.C. § 119 12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of: 1. Certified copies of the priority documents have been received. 2. Certified copies of the priority documents have been received in Application No 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received.	4a) Of the above claim(s) <u>268-292</u> is/are withdrawn from consideration.						
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DETAILED ACTION

Claims 224, 228, 229, 232-292 are pending. Claims 224, 228, 229, 232-267 are considered on the merits. Claims 268-292 are withdrawn from consideration as being drawn to a non-elected invention.

Once again applicants are requested to cancel the non-elected claims as instructed in the final office action of 12/17/03 in order to prepare the application in the event of allowance.

The text of those sections of Title 35, U.S. Code not included in this action can be found in a prior Office action.

Claim Rejections - 35 USC § 103

Claims 224, 228, 229, 232-245, 248-252, 255, 258-266 remain rejected under 35 U.S.C. 103(a) as being unpatentable over WO 88/05261 [N].

The claims are directed to a method of maintaining and restoring the viability of an organ subjected to ischemia comprising:

perfusing the organ with a first fluid at a first temperature to maintain/restore pre-ischemia energy levels in the organ, and

perfusing the organ with a second fluid containing substantially no oxygen at a second temperature to store/transport the organ, whereby the second temperature is lower than the first temperature, wherein the first temperature is from about 12°C to about 24°C.

The references are relied upon as explained below.

WO 88/05261 discloses in Example 3, a first normothermic (37°C) perfusion of a FC-43 emulsion at a pressure of 40-120mm Hg followed by a second perfusion with a hypothermic (4-6°C) electrolyte solution at a pressure of <20mm Hg. The perfusate is circulated through a pH sensor module (page

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25, l. 20) which is a viability marker or indicative of the organ's viability according to the specification at page 15, l. 28. The perfusate is recycled through a recirculation loop which contains a filter (Fig 1, 16). The recycled perfusate is debubbled and oxygenated and pH regulated prior to returning to the organ. The perfusion may be continuous during storage. The organ is first removed from the subject and cooled in a saline/icewater bath (page 22, l. 10) and perfused with a cardioplegia solution prior to perfusing with the first and second solutions. The organ is stored after the second perfusion in a chamber with includes a housing and an organ supporting surface which allows effluent to pass through, the housing includes openings.

The FC-43 emulsion is an oxygen carrying solution which contains dextrose, both of which function to maintain energy levels in organs (See Ingwall *et al.* [U]). The solution contains oxygen and the partial pressure of oxygen is monitored during the perfusion process. Thus the solution is considered to have a viability marker as disclosed on page 6, l. 19 of the specification.

The electrolyte solution contains mannitol which is an antioxidant (See Burdon et al. [V]).

Insofar as the process relies on the use of components of the solutions which instead of being characterized by technical features suitable for the identification of a solution composition, is imprecisely defined by means of functional features which merely recite the desired result to be achieved, the subject matter is considered to be anticipated or made obvious by the disclosures of the prior art.

Portable is a term without a reference point. Everything is portable if a large enough moving force is applied. Thus, the chamber (1) of the prior art which holds the artificial pericardial sack (2) and has inlets (51) and outlets (16) connected to the organ, can be portaged and thus the chamber is considered to be portable. That the applicants have termed the pericardial sack (2) a

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"cassette" is the prerogative of the inventor and as it is lacking in structural elements is of little patentable weight. Please note the artificial pericardial sack may be thrown away after use if wished, thus it may be termed "disposable". The container is capable of maintaining the organ at a temperature of 10°C at least for a short period of time because the heart has been cooled. No length of time of the "maintaining" limitation is seen. Maintaining can be for one minute, one hour, or less.

The reference specifically lacks the temperature limitation for the first solution perfusion which is now about 12-24°C. Instead the first perfusion is performed at 37°C.

With regard to the limitation of the first temperature being about 12°C to about 24°C, WO 88/05261 teaches the first temperature of 37±1°C. The instant specification teaches a range of 10–38°C for the first solution. Thus, the new range limitation is merely an optimization of ranges and is still considered to be obvious over the prior art in the absence of criticality. See MPEP 2144.05 ll.A. Also, please note on page 21, lines 15–20, that at temperatures below 20°C, the mitochondria are unable to produce energy. Use of temperatures below 20°C, therefore would not be expected to significantly contribute to restoring energy levels in the organ. Evidence to the contrary might provide unexpected results.

Claim 255 is directed to transplanting the organ into the recipient while the organ remains at the second temperature which is the at temperature that it last was perfused.

While WO 88/05261 does not teach any specific temperature that the heart must be during transplantation, in the absence of evidence to the contrary, one of skill in the art may chose to transplant a cold organ or a warm one as desired.

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Claims 246 and 247 remain rejected under 35 U.S.C. 103(a) as being unpatentable over WO 88/05261 [N] in view of WO 96/29864 [O]

The claims are further directed to the use of a pressure source incapable of providing pressures greater than 100 mmHg or 40 mmHg.

WO 88/05261 discloses using 40–120 mm Hg pressures in normothermic perfusion and less than 20 mm Hg pressures in hypothermic perfusion. The system is equipped with a pressure release control system which is programmable so as not to exceed a preestablished limit (page 14, ls. 1–10). Fig 4 shows the control systems. The reference lacks the disclosure of using a setting of 100 mmHg which cannot be exceeded.

WO 96/29864 discloses an apparatus used for normothermic perfusing of organs with a pressure maximum of 90 mmHg (page 23).

One of skill in the art may set the pressure in the apparatus of WO 88/05261 so that the fluid perfusion pressure can not exceed 90 mmHg as taught by WO 96/29864 because the apparatus of WO 88/05261 has variable settings which once established according to the desire of the operator will not be exceeded. One of skill in the art would be expected to able to establish maximal settings of 90 mmHg using the apparatus of WO 88/05261 and the maximal pressure settings as disclosed in WO 96/29864 in the absence of evidence to the contrary.

Clearly the apparatus is capable of operating at pressures no greater than 20mmHg also because this is the setting taught for hypothermic perfusion.

Claims 253 and 254 remain rejected under 35 U.S.C. 103(a) as being unpatentable over WO 88/05261 [N] in view of Chambers *et al.* [W].

The claims are directed to the use of antioxidants in the flush solution.

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WO 88/05261 is relied upon as discussed above and lacks the specific teaching of the use of antioxidants in the flush/cardioplegic solution.

Chambers et al. disclose that the addition of various antioxidants to cardioplegic solutions improves organ viability.

The addition of an antioxidant to the flush, cardioplegic solution of WO 88/05261 (p. 22, l. 6) would have been obvious when taken with Chambers *et al.* who teach the improved viability of a heart when the flush/cardioplegic solution incorporates antioxidants.

Claim 257 remains rejected under 35 U.S.C. 103(a) as being unpatentable over WO 88/05261 [N] and Ingwall *et al.* [U] or WO 97/43899 [P].

The claim is direct to perfusing the organ with the first fluid at the first temperature prior to transplantation.

WO 88/05261 teaches at page 23, that it is possible to raise the temperature of the heart if there is ATP depletion prior to transplantation.

WO 97/43899 disclose that warming a heart (page 10, I. 23) and perfusing with a solution containing substrates such as glucose (page 10, I. 33) and oxygen (page 11, I. 3) reestablishes oxidative metabolism, which means that ATP levels are established (abstract and Fig. 1).

Inwall et al. teaches that ATP levels can be restored in heart when perfusion with a solution containing oxygen and glucose (abstract).

One of skill in the art, given the teaching that one should raise the temperature of the heart if there is ATP depletion (WO 88/05261), would perfuse with the first fluid at the first temperature because warm perfusion of the heart with oxygen and glucose is known to raise endogenous ATP levels and improve viability of the organ as taught by Ingwall *et al.* or WO 97/43899 [M].

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Claims 259-266 remain rejected under 35 U.S.C. 103(a) as being unpatentable over WO 88/05261 [N] in view of US 5,586,438 [A].

The claims are directed to a method of perfusing an organ in a portable perfusion unit capable of maintaining the organ at a temperature of 10°C or less. Other claims are directed to storing the organ after perfusion with the first medical fluid at the first temperature or with the second medical fluid at the second temperature.

The references are relied upon as explained below.

WO 88/05261 has been discussed above and lacks specific mention of portability.

US 5,586,438 discloses an apparatus for transporting and preserving organs (Fig. 3). It comprises a housing and an organ container. The apparatus can be cooled by ice or other thermal buffers or by the expansion of compressed gas. The preferred temperature of storage/transportation is 6°C (col. 9, I. 29). While the organ container is not disclosed as being disposable, anything in the absence of structural elements can be disposed of. The organ container is removable from the housing and thus, can be thrown away separate from the housing.

The use of the perfusion regimen of WO 88/05261 in the apparatus of US 5,586,438 would have been obvious because one of skill in the art may choose any perfusion apparatus in the art that will perform a specific perfusion regimen. The apparatus of US 5,586,438 is capable of performing the perfusion regimen taught in WO 88/05261 and is used for perfusion and transportation of organs for transplantation.

Further, it is well within the purview of one of skill in the art to store/transport the organ after the perfusion of any solution known in the art at any temperature used in the art in the absence of any evidence of criticality.

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Claim 267 remains rejected under 35 U.S.C. 103(a) as being unpatentable over WO 88/05261 [N] and US 5,586,438 [A] as applied to the claims above, and further in view of US 5,450,329 [B].

The claims are further directed to the use of a GPS to monitor the location of the organ.

WO 88/05261 and US 5,586,438 are relied upon as discussed above. The references lack mention of the use of GPS to track the organ.

US 5,450,329 discloses the use of GPS to track a vehicle.

The use of GPS to track a vehicle which is carrying the portable perfusion unit of US 5,450,329 which contains an organ perfused with the regimen of WO 88/05261 and is intended for transplantation is well within the purview of one of skill in the art because such tracking devices are known in the art.

One of ordinary skill in the art would have been motivated at the time of invention to make these substitution in order to obtain the results as suggested by the references with a reasonable expectation of success. The claimed subject matter fails to patentably distinguish over the state of the art as represented by the cited references. Therefore, the claims are properly rejected under 35 U.S.C. § 103.

No critical or novel element is seen in the claims. It appears that all elements are known in the art of transplantation.

Response to Arguments

Applicants' arguments filed 10/15/04 have been fully considered but they are not persuasive.

Applicants argue that the temperature range of 12-24°C is critical because it allows some specific metabolic results to come to pass while

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preventing others. Applicants further argree that mitochondial activity is reduced or arrested from 12-24°C, but that the glucose-pyruvate-lactate pathway is active and is producing ATP in the cytosol. Also mentioned is the assertion that the killing rates of cells exposed to reduced temperatures changes the slope of an Arrhenious plot in the region of 8-10C and on this basis there "could be" distinct advantages of storing organs above the transistion temperature.

While this is an interesting discussion and may point the direction for evidence to be presented, this is merely an argument of counsel without any comparative evidence with the method of the prior art. Particularly since the specification states on page 21, that that the first perfusion temperature may be normothermic and the first perfusion solution may include oxygen to enable the mitochondria to replenish the ATP levels. The thrust of the disclosure in the specification appears to be a first normothermic or near normothermic perfusion with an oxygen supplying first solution to allow the "maintaining or restoring viability", ATP levels by mitochondrial metabolism, and then a hypothermic perfusion with a second solution that does not supply oxygen to prevent degradation of the cells. No evidence or even mention of the criticality of a first perfusion in the range of 12-24°C appears in the specification, merely a mention of "perferably, the normothermic temperatures are from about 12-24°C, but higher or lower temperatures can be used as desired or necessary". On page 26, normotheric or near normotheric are defined in a range of 10-38°C which is an overlapping temperature with the prior art. As no evidence of this criticality has been presented by way of declaration in a comparative example with the method of the prior cited art, applicants arguments are interesting, but unpersuasive in overcoming the prior art disclosure.

Clear evidence that a first perfusion with an oxygen carrying solution in the range of 10-24° C is superior to a similar perfusion around 37°C for the maintaining and/or restoring the viability of an organ sujected to a period of ischemia or hypoxia, might advance applicants' prosecution.

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Conclusion

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Sandra Saucier whose telephone number is (571) 272-0922. The examiner can normally be reached on Monday, Tuesday, Wednesday.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, M. Wityshyn can be reached on (571) 272-0926. The fax phone number for the organization where this application or proceeding is assigned is 703-872-9306.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free).

Sandra Saucier

Primary Examiner

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December 13, 2004